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(71) Applicant: **WESTINGHOUSE ELECTRIC CORPORATION**
Westinghouse Building Gateway Center
Pittsburgh Pennsylvania 15222(US)

(72) Inventor: **Harrison, Don Edward**
3809 Edinburg Drive
Murrysville Pennsylvania(US)

(72) Inventor: **Pope, James Michael**
4427 Helmar Drive
Monroeville Pennsylvania(US)

(72) Inventor: **Wood, Susan**
2374 Collins Road
Pittsburg Pennsylvania(US)

(74) Representative: **van Berlyn, Ronald**
23 Centre Heights
London NW3 6JG(GB)

(54) **Method of immobilizing nuclear waste in glass.**

(57) Disclosed is a method of immobilizing nuclear waste in glass. A composition is prepared of 60 to 100% of a hydrolyzed glass-forming silicon compound and up to about 40% of a glass-forming aluminum compound. About 1 to about 50% liquid nuclear waste and up to about 10% solid nuclear waste is mixed into the composition. The composition is heated to about 200 to about 500°C to drive off water and organics, with the resulting vitreous product totally containing the nuclear waste. Finally, this product can be sintered at about 800 to about 900°C to reduce porosity, or worm pressed into block form.

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METHOD OF IMMOBILIZING NUCLEAR WASTE

This invention relates to a method of immobilizing nuclear waste.

Reprocessing of either spent nuclear fuel or weapons material results in liquid waste which must be reduced in volume and consolidated to permit safe disposal. The current practice is to dehydrate the liquid waste by heating, then to consolidate the residue by either calcination or vitrification at high temperatures. In the past, defense waste was neutralized in order to precipitate metallic hydroxides. This product can be converted into a vitreous waste form using conventional glass forming technology.

The ultimate suitability of vitreous waste forms is suggested by the durability of rhyolytic obsidian and tektite natural glasses during millions of years in a variety of geologic environments. Unfortunately, these chemically durable, high-silica glasses pose problems as a practical solid-waste form when made using conventional continuous vitrification processes. Because of the high fluxing temperatures (~1350°C) required, additional off-gassing scrubbing capacity or other absorbent procedures are needed to deal with the volatilization losses of radionuclides such as iodine, cesium, and ruthenium. The high fluxing temperatures also shorten furnace life, and can create problems with the materials into which the molten glass is cast, such as the sensitization of stainless steel to stress corrosion cracking. As a consequence

of these limitations, most nuclear waste glass formulations have substantially lower silica content than either natural obsidians, nepheline syenite, or commercial "Pyrex" glasses. Less silica or alumina and more fluxing agent (e.g., Na_2O , K_2O or B_2O_3) lowers the glass working temperature (to 1000-1200°C for most waste glasses) and raises the waste loading capacity. However, this also results in lower chemical durability in most aqueous environments and, particularly for borosilicate compositions, in less resistance to devitrification.

We have now discovered that the formation of aluminosilicate glasses by chemical polymerization can effectively contain nuclear waste. The process of this invention avoids the volatilization losses that occur with conventional glass-forming processes because the temperatures used in the process of this invention are relatively low. The invention immobilizes the nuclear waste in a highly leach resistant glass which could not be formed by prior processes except at very high temperatures.

According to the present invention, a method of immobilizing nuclear waste is characterized by (A) preparing a composition which comprises: (1) from 60% to about 100% by weight, calculated as SiO_2 , of a hydrolyzed silicon compound having the general formula $\text{SiR}_m(\text{OR}')_n\text{X}_p$ or $\text{Si}(\text{OSiR})_4$ where each R is independently selected from alkyl to C_{10} and alkenyl to C_{10} , each R' is independently selected from R and aryl, each X is independently selected from chlorine and bromine, m is 0 to 3, n is 0 to 4, p is 0 to 1, and $m + n + p$ equals 4; (2) up to about 40% by weight, calculated as Al_2O_3 , of an aluminum compound having the general formula $\text{AlR}'_q(\text{OR})_r\text{X}_s$ or $\text{Mg}(\text{Al}(\text{OR})_4)_2$, where each R is independently selected from alkyl to C_{10} and alkenyl to C_{10} , each R' is independently selected from R or aryl, q is 0 to 3, r is 0 to 3, s is 0 to 1, and $q + r + s$ equals 3; (B) mixing from 1 to 50%, based on total weight, of said nuclear waste in liquid form into said composition; (C) mixing up to about 10%,

based on total weight, of said nuclear waste in solid form into said composition; and (D) heating said composition containing said nuclear waste at from 200 to 500°C to drive off water and organics.

5 The $\text{SiR}_m(\text{OR}')_n\text{X}_p$ compounds are preferred as those compounds are more available, easier to handle and more compatible. The R' group is preferably alkyl to C_4 with $n = 4$ because alkoxides are the most suitable starting compounds.

10 Appropriate compounds which fall within the scope of the general formula include

trimethylethoxysilane, $(\text{CH}_3)_3\text{Si}(\text{OC}_2\text{H}_5)$

ethyltriethoxysilane $\text{C}_2\text{H}_5\text{Si}(\text{OC}_2\text{H}_5)_3$

tetrapropoxysilane $\text{Si}(\text{OC}_3\text{H}_7)_4$

15 tetraethylorthosilicate $\text{Si}(\text{OC}_2\text{H}_5)_4$

tetratriethylsiloxysilane $\text{Si}[\text{OSi}(\text{CH}_3)_2\text{C}_2\text{H}_5]_4$

triethylchlorosilane $(\text{C}_2\text{H}_5)_3\text{SiCl}$

vinyltriphenoxysilane $\text{CH}_2:\text{CHSi}(\text{OC}_6\text{H}_5)_3$.

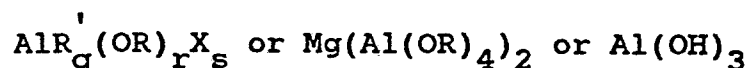
20 The preferred silicon compound is tetraethylorthosilicate because it is relatively inexpensive, readily available, stable, and easy to handle. The above compounds are partially hydrolyzed with water in alcohol. It is preferable to partially hydrolyze the silicon compound prior to mixing it with the other components because its rate of

25 hydrolysis is slower and precipitation may occur if hydrolysis is done after mixing. It is preferable to use the same alcohol that is formed during subsequent polymerization so that two alcohols need not be separated. A suitable molar ratio of the silicon compound to the al-

30 cohol is about 0.2 to about 2. A suitable molar ratio of the silicon compound to the water used in hydrolysis is from 0.1 to 5. In addition, it is sometimes helpful to

add up to about 6 drops of concentrated nitric acid per mole of water to aid in hydrolyzation. After the water is added to the silicon compound the compound is permitted to sit for several hours to permit hydrolyzation to occur.

- 5 The aluminum compounds which are suitable for use in this invention have the general formula:



where each R' is independently selected from R and aryl, q is 0 to 3, r is 0 to 3, s is 0 to 1, and $q + r + s = 3$.

- 10 The $\text{AlR}'_q(\text{OR})_r\text{X}_s$ compounds, where r is 3 and R is alkyl to C_4 , are preferred as they are the most stable and available and are easiest to handle. The R group in the aluminum compound need not be the same R group that is in the silicon compound.

- 15 Suitable compounds which fall within the scope of the general formula include:

	trimethylaluminum	$\text{Al}(\text{CH}_3)_3$
	triethylaluminum	$\text{Al}(\text{C}_2\text{H}_5)_3$
	triethoxyaluminum	$\text{Al}(\text{OC}_2\text{H}_5)_3$
20	aluminum isoproponate	$\text{Al}(\text{OC}_3\text{H}_7)_3$
	aluminum secondary butoxide	$\text{Al}(\text{OC}_4\text{H}_9)_3$
	triphenyl aluminum	$\text{Al}(\text{C}_6\text{H}_5)_3$
	aluminum magnesium ethoxide	$\text{Mg}[\text{Al}(\text{OC}_2\text{H}_5)_4]_2$
	diethylaluminum chloride	$(\text{C}_2\text{H}_5)_2\text{AlCl}$

- 25 The preferred aluminum compounds is aluminum secondary butoxide because it is stable, available, and does not require special handling. The aluminum compound (other than the hydroxide) is preferably hydrolyzed before it is added to the silicon compound because the mixture

will then act compatibly as a single compound and inhomogeneities will be avoided. The molar ratio of the aluminum compound to the water used to hydrolyze it can range from 0.0007 to 0.03. The water should be hot (i.e.,
5 between 70 and 100°C, and preferably between 80 and 90°C) to facilitate proper hydrolyzation. In addition, it may be desirable to use from 0.03 to 0.1 moles of 1 molar nitric acid per mole of $AlO(OH)$, which is the desired product of the hydrolyzation, to aid in its peptization.
10 After the addition of the water, the compound is permitted to set for at least several hours at from 80 to 90°C to permit proper hydrolyzation and peptization to occur.

After the silicon compound and the aluminum compound have been separately hydrolyzed they are mixed to
15 prepare the composition. The composition may include from 60 to 100% by weight of the silicon compound calculated as SiO_2 and based on the total weight of $SiO_2 + Al_2O_3$ and up to about 40% by weight of the aluminum compound calculated as Al_2O_3 based on the total weight of $SiO_2 + Al_2O_3$.
20 Preferably, the composition comprises from 70% to 90% by weight of the silicon compound calculated as SiO_2 and from 10% to 30% of the aluminum compound calculated as Al_2O_3 , because more than about 30% of the aluminum compound may make the composition more difficult to warm press. At
25 less than about 10% of the aluminum compound the durability of the glass may suffer.

The composition can immobilize both solid nuclear waste and an aqueous solution of nuclear waste. The dissolved nuclear waste is usually nitrate solutions of
30 various metals including iron, uranium, nickel, magnesium, calcium, zirconium, plutonium, chromium, cobalt, strontium, ruthenium, copper, cesium, sodium, cerium, americium, niobium, thorium, and curium. Depending on the species present, it may be preferable to adjust the pH of the nuclear waste solution with a hydroxide so that is approxi-
35 mates the pH of the glass composition. The dissolved nuclear waste can contain from about 5% dissolved solids

to saturated, and a typical solution of nuclear waste may have from 10% to 30% solids in solution. For example, a typical nuclear waste is up to about 15% by weight nitrate and up to about 85% by weight water. Up to about 50%
5 based on the total weight of the waste plus the glass composition can be nuclear waste in liquid form.

Solid nuclear waste can also be added to the glass composition. Solid nuclear waste generally consists of the hydrated oxides and hydroxides, and possibly sulfates, phosphates, nitrites or other salts of the metals
10 listed above. Up to about 10% based on the total weight of the nuclear waste and the composition may consist of solid nuclear waste.

The nuclear waste material is added to the glass composition with stirring and the mixture is dried. The drying, which polymerizes the silicon and aluminum oxides, may begin at room temperature and extend to about 150°C at a rate of temperature increase of from 1°C to 10°C per minute. Between 150°C and 200°C the mixture may be heated
15 more rapidly (e.g., at a rate of temperature increase of from 10°C to 50°C per minute) in order to more effectively drive off the carbon. Finally, between 200°C and 500°C the mixture is again heated at the slower rate of temperature increase of from 1°C to 10°C per minute in order to
20 remove the remaining water of hydration and any organics which may be present.

The resultant 500°C product is vitreous granules, about 1-10 mm in diameter, which effectively contain the nuclear waste. This containment is generally by
30 complete dissolution in glass, although encapsulation in the sense that certain few insoluble species are totally surrounded by the glass may also occur. The granules typically have a high surface area, although their durability and stability do not appear to be adversely affected. Nevertheless, it may be desirable to further
35 process the granules. For example, sintering at from 800°C to 900°C for up to about 10 hours will reduce the

surface area of the granules from 500 m²/g to less than approximately 10 m²/g.

To prepare a solid block of contained and immobilized nuclear waste the waste-glass granules are warm pressed at from 350°C to 600°C using from 30,000 to 150,000 psi, depending on the temperature. The higher the temperature, the lower is the pressure that will be needed, and the lower the temperature is, the higher the pressure will need to be in order to produce a solid block. After about one half hour of warm pressing a solid block of the immobilized waste is produced. The following example further illustrates this invention.

The invention will now be illustrated with reference to the following Example:

15

EXAMPLE

The following compounds were added in sequence at room temperature.

90 grams of pure ethyl alcohol

20

9 grams of deionized water (1 mole H₂O/mole tetraethylorthosilicate)

1 drop concentrated (7.45 M) HNO₃

104 grams tetraethylorthosilicate

The composition was stirred for 15 minutes, covered tightly and allowed to age at room temperature for 16 hours. An aluminum monohydroxide composition was prepared by heating 162 grams deionized water to 85°C, adding 16 grams of aluminum secondary butoxide while stirring, and adding 4 cubic centimeters of 1 M HNO₃ (moles acid/moles aluminum equals 0.06). The composition was stirred for 15 minutes, covered and allowed to age at 85°C for 16 hours. The aluminum monohydroxide composition was then added to the siloxane composition at room temperature with stirring.

A surrogate liquid waste composition was prepared by dissolving the following nitrates in 10 cc deionized water.

3.1990 grams $\text{Fe}(\text{NO}_3)_3 \cdot 9\text{H}_2\text{O}$

0.9330 grams $\text{UO}_2(\text{NO}_3)_2 \cdot 6\text{H}_2\text{O}$

1.0684 grams $\text{Sr}(\text{NO}_3)_2$

1.6346 grams NaNO_3

5 Within 2-3 minutes after the siloxane and aluminum monohydroxide compositions were mixed, the surrogate liquid waste was added in the order listed while stirring at room temperature.

10 Alternatively, up to about 2% by weight of a surrogate solid waste (apatite) was added to the room temperature siloxane-aluminum monohydroxide mixture while stirring. The mixture was stirred and heat was applied at about 125 to 150°C until a gel formed and was subsequently dried.

15 Generally, the volume reduction was about 33% to reach the gelatinous state and approximately an additional 33 vol% shrinkage occurred in obtaining a dried material. The total volume reduction was less with the solid waste loading, being about 50% at a 10% waste level. Using a
20 quartz tray, a fairly thin bed of material was heated to 500°C in air. The heating rate was about 1°C per minute to 150°C, followed by rapid heating of about 10°C per minute to 225°C, then about 1°C per minute to 500 or 850°C. The material was held at 500°C for 16 hours. The
25 result was a totally amorphous granular material having a grain size of about 1 to 10 mm.

30 A second surrogate solid waste was prepared and tested in the same manner as the apatite. The second surrogate waste form simulated the analyzed composition of an actual sample of nuclear waste and had the following composition.

	$\text{Fe}_2(\text{SO}_4)_3$	54.5 wt%
	$\text{Al}_2(\text{SO}_4)_3$	4.9 wt%
	MnSO_4	3.3 wt%
	$\text{UO}_2(\text{NO}_3)_2$	15.3 wt%
5	Na_2PO_4	4.9 wt%
	$\text{Sr}(\text{NO}_3)_2$	2.8 wt%
	CaSO_4	3.8 wt%
	NiPO_4	10.5 wt%

10 The amounts of this waste added to the mixed gel derivatives and also the gel were 1.0, 5.0 and 10.0 wt% total metal with respect to the Si plus Al. The following table gives the results of leach tests on these samples.

Conditions During Leach Tests	Species Tested in Emission Spectrographic Analysis ($\mu\text{g/ml}$)									Leach Rate (total $\text{g/cm}^2/\text{day}$)
	Al	Si	Fe	U	Sr	Na	Ca	P	F	
<u>Glass Containing Waste</u> Test Water ¹ Static, R.T. H_2O , 2 wk ² Static, R.T. H_2O , 8 wk ³ Soxhlet, 100°C H_2O , 8 wk	<0.1 <0.5 1.1 2.8	3.6 9.5 168 947	<0.1 <0.5 0.43 1.8	<0.1 <0.5 2.0 7.3	<0.1 <0.1 0.23 0.39	<0.1 1.6 83 84				3.98×10^{-11} 2.71×10^{-10} 1.05×10^{-10}
<u>Glass Containing 1% Apatite</u> Test Water ⁴ Soxhlet, 100°C H_2O , 6 wk	0.46	38				0.86	1.4	0.79	0.30	1.67×10^{-10}

Other details concerning samples:

- | Details concerning samples | Surface area of sample, m^2/g |
|--|--|
| 1 3780 ml H_2O , 14.50g of sample, | 421 m^2/g surface area of sample. |
| 2 3700 ml H_2O , 14.50g of sample, | 421 m^2/g surface area of sample. |
| 3 450 ml H_2O , 18.95g of sample, | 421 m^2/g surface area of sample. |
| 4 450 ml H_2O , 35.23g of sample, | 6.95 m^2/g surface area of sample. |

Water pH at the conclusion of the experiments was 6.2-6.3 in all cases.

CLAIMS:

1. A method of immobilizing nuclear waste characterized by (A) preparing a composition which comprises: (1) from 60% to about 100% by weight, calculated as SiO_2 , of a hydrolyzed silicon compound having the general formula $\text{SiR}_m(\text{OR}')_n\text{X}_p$ or $\text{Si}(\text{OSiR})_4$ where each R is independently selected from alkyl to C_{10} and alkenyl to C_{10} , each R' is independently selected from R and aryl, each X is independently selected from chlorine and bromine, m is 0 to 3, n is 0 to 4, p is 0 to 1, and $m + n + p$ equals 4; (2) up to about 40% by weight, calculated as Al_2O_3 , of an aluminum compound having the general formula $\text{AlR}'_q(\text{OR})_r\text{X}_s$ or $\text{Mg}(\text{Al}(\text{OR})_4)_2$, where each R is independently selected from alkyl to C_{10} and alkenyl to C_{10} , each R' is independently selected from R or aryl, q is 0 to 3, r is 0 to 3, s is 0 to 1, and $q + r + s$ equals 3; (B) mixing from 1 to 50%, based on total weight, of said nuclear waste in liquid form into said composition; (C) mixing up to about 10%, based on total weight, of said nuclear waste in solid form into said composition; and (D) heating said composition containing said nuclear waste at from 200 to 500°C to drive off water and organics.

2. A method according to claim 1, characterized by including, after driving off water and organics, the additional step of sintering the composition at from 800 to 900°C.

3. A method according to claim 1 or 2, characterized by the last step of warm pressing said composition at from 350 to 600°C at from 30,000 to 150,000 psi.

4. A method according to claim 1, 2 or 3, characterized in that the nuclear waste is about 5% to saturated with solids and comprises about up to about 15% nitrate, up to 85% water, and up to about 10% undissolved solids.

5. A method according to claim 1, 2, 3 or 4, characterized in that the silicon compound has the general formula $\text{SiR}_m(\text{OR}')_n\text{X}_p$ where R' is alkyl to C_4 and $n = 4$ and the aluminum compound has the general formula $\text{AlR}'_q(\text{OR})_r\text{X}_s$ where R is alkyl to C_4 and r is 3.

6. A method according to claim 5 characterized in that the silicon compound is tetraethylorthosilicate and the aluminum compound is aluminum secondary butoxide.

7. A method according to any of the preceding claims, characterized in that the silicon compound is hydrolyzed in alcohol at a molar ratio of silicon compound to alcohol of from 0.2 to 2, with water at a molar ratio of silicon compound to water of from 0.1 to 5.

8. A method according to any of the preceding claims, characterized in that the composition comprises from 70 to 90% of the silicon compound and from 10 to 30% of the aluminum compound.